

1    Rifting under steam – how rift magmatism triggers  
2    methane venting from sedimentary basins

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**ABSTRACT**

During opening of a new ocean magma intrudes into the surrounding sedimentary basins. Heat provided by the intrusions matures the host rock creating metamorphic aureoles potentially releasing large amounts of hydrocarbons. These hydrocarbons may migrate to the seafloor in hydrothermal vent complexes in sufficient volumes to trigger global warming, e.g. during the Paleocene Eocene Thermal Maximum (PETM). Mound structures at the top of buried hydrothermal vent complexes observed in seismic data off Norway were previously interpreted as mud volcanoes and the amount of released hydrocarbon was estimated based on this interpretation. Here, we present new geophysical and geochemical data from the Gulf of California suggesting that such mound structures could in fact be edifices constructed by the growth of black-smoker type chimneys rather than mud volcanoes. We have evidence for two buried and one active hydrothermal vent system outside the rift axis. The vent releases several hundred degrees Celsius hot fluids containing abundant methane, mid-ocean-ridge-basalt (MORB)-type helium, and precipitating solids up to 300 m high into the water column. Our observations challenge the idea that methane is emitted slowly from rift-related vents. The association of large amounts of methane with hydrothermal fluids that enter the water column at high pressure and temperature provides an efficient mechanism to transport hydrocarbons into the water column and atmosphere, lending support to the hypothesis that rapid climate change such as during the PETM can be triggered by magmatic intrusions into organic-rich sedimentary basins.

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## 52 INTRODUCTION

53 When testing climate models by hind casts for past prominent warming events it has  
 54 become clear that significant drivers are missing (Panchuk et al., 2008; Carozza et  
 55 al., 2011). Most striking, the reason for a 5-6°C global warming during the PETM  
 56 remains elusive, although it is clear that more than 2000 Gt of carbon must have  
 57 entered the atmosphere within a relatively short time span of a few thousand years  
 58 (Dickens et al., 1995; Zachos et al., 2001). Light carbon isotope ratios were  
 59 interpreted to suggest that large amounts of biogenic carbon were released through  
 60 run-away hydrate dissociation events (Kennett et al., 2000). But recent studies call  
 61 this interpretation in question (Dickens, 2011; Biastoch et al., 2011). Most  
 62 alternative explanations are also marred by contradictions with available data  
 63 (Higgins and Schrag, 2006). This includes the hypothesis of carbon mobilization by  
 64 magmatic intrusions into carbon-rich continental margin sediments during the  
 65 opening of the North Atlantic (Svensen et al., 2004), because it is difficult to  
 66 understand how this process could have injected large amounts of carbon into the  
 67 atmosphere within a short time.

68

## 69 THE ACTIVE HYDROTHERMAL MOUND IN THE GUAYMAS BASIN

70 New data acquired in the Guaymas Basin (Fig. 1, S1) support the ‘tectonic-magmatic’  
 71 explanation for warming during the PETM. Our data show a 1050 m-wide and 100 m-  
 72 high mound located above thick sediments approximately 1 km southeast of the  
 73 northern rift axis of the Guaymas Basin that is emitting carbon-rich hydrothermal  
 74 fluids (Fig. 2a). The mound and most of the southern flank of the Guaymas rift valley

are underlain by an approximately 100 m-thick chaotic seismic facies (50 to 150 ms two-way-travel time (TWT)) that is part of a widespread mass transport deposit. In the area of the mound another chaotic and low seismic amplitude seismic facies continues further downward to at least 2900 ms TWT or approximately 200-250 metres below seafloor. We interpret this facies as a conduit for the ascending hydrothermal fluids. At this depth high seismic amplitude reflections extend southward from below the rift axis into the basin. These reflections are likely caused by magmatic intrusions which occur throughout the Guaymas Basin (Lizarralde et al., 2010). The bathymetric data (Fig. 1) also show the trace of a NE-SW striking normal fault that stretches to the mound and is dipping westward (Fig. S2). As it strikes in the same direction as the elongate mound axis it is probable that the shape of the mound is controlled by this fault. An approximately 20 m-thick chaotic seismic facies immediately below the base of the mound may indicate that hydrothermal seepage was initially occurring over a broader area and became more focused once fluid pathways had formed. The up to 30°-steep flanks of the mound suggest a high internal angle of repose of the mound-forming material.

A 2.2 km-long profile of ten heat flow stations across the vent field and a 0.4 km-long profile of five stations along the mound consistently show heat flow values larger than 0.3 W/m<sup>2</sup>. The maximum heat flow on top of the mound is heterogeneous with values exceeding 10 W/m<sup>2</sup>, whereas values below 3W/m<sup>2</sup> were measured only 100 m away from the center (Supplementary Table S1). This heat flow pattern is similar to the heat flow distribution at sediment-free mid ocean ridges (Rona et al., 1996). The strong lateral heat flow variations indicate focused heat transport by hydrothermal fluids within narrow fluid pathways.

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101 Samples recovered from the top of the mound show porous Fe-rich sulfides (Fig. S3)  
102 consisting predominantly of a boxwork of pyrrhotite crystals with minor pyrite, and  
103 marcasite, and rare Zn- and Cu-sulfides (e.g. sphalerite, chalcopyrite, isocubanite).  
104 These are accompanied by a small fraction of non-sulfide minerals (e.g., carbonates,  
105 opal-A and secondary Fe-oxyhydroxides). The mineralogy makes the samples from  
106 the northern rift axis similar to the massive sulfides recovered from the southern  
107 Guaymas Basin (Koski et al., 1985; Peter and Scott, 1988). However, they lack the  
108 abundant petroleum found in the southern rift samples. In analogy to black smoker  
109 deposits from Escanaba Trough the observed mineral assemblage suggests formation  
110 temperatures in the range of 270-330°C (Zierenberg et al., 1993).

111

## 112 **TIMING OF HYDROTHERMAL ACTIVITY**

113 A 5m-long gravity core was taken ~500 m away from the hydrothermal vent field  
114 (Fig. 3). The upper 4 m of the core consist of organic-rich, hemipelagic diatomaceous  
115 clay that is typical for most of Guaymas Basin (Damm et al., 1984). By contrast, the  
116 lowermost 1 m of core consists of coarse-grained hydrothermal deposits intercalated  
117 with clay lenses suggesting the mound to consist of hydrothermal deposits. Given Pb-  
118 210-based sedimentation rates of 0.74 m/kyr inside the hydrothermal vent field  
119 (Station 40) and of 0.79 m/kyr just outside the hydrothermal vent field along the  
120 Northern Guaymas rift axis (Station 66), the hydrothermal deposits are likely 5 to 6  
121 kyr old. This is a minimum age estimate as older hydrothermal material likely exists  
122 below and towards the centre of the vent complex. However, we note that sill  
123 intrusions quickly cool off after emplacement (Jamtveit et al., 2004) and even thick

intrusions can only sustain hydrothermal systems for a few kyr. Thus, we infer that the mound structure must have formed during the last 5-10 kyr.

## ORIGIN OF THE HYDROTHERMAL FLUIDS

We collected sea water in the vicinity of the hydrothermal field (Fig. 3) with Niskin bottles and measured the dissolved gases with an adapted membrane-inlet mass-spectrometer (MIMS (Mächler et al., 2012)). The water is strongly enriched in helium (He) relative to the atmospheric equilibrium conditions (Fig. S5). At the flanks and the bottom of the rift valley He concentrations are similar to the concentrations found in the southern part of the Guaymas Basin (Lupton, 1979) whereas He in the water just above the active smokers is supersaturated by more than a factor of 4 (Fig. S6). This enrichment indicates that the vent injects He directly into the water column. The injected He is strongly enriched in  $^3\text{He}$ . The  $^3\text{He}/^4\text{He}$  ratio ( $10.8 \cdot 10^{-6}$ , Fig. S6, Tab. S3) agrees with that of excess He from the southern part of the Guaymas Basin (Lupton, 1979) pointing to the same general MORB source, and confirms that water circulation in the Guaymas Basin distributes He-rich water from the black smoker region throughout the basin (Marinone, 2003).

Dissolved concentration of light hydrocarbons determined in water sampled at the smoker field (Fig. 3) by Niskin bottles are highly enriched (factor  $\geq 10^5$ ) compared to bottom water concentrations (Tab. S2). Calculated end member concentrations (i.e.  $\text{CH}_4 = 6.5 \text{ mM}$ ; Fig. S7) are similar to that of vent fluids measured in the Southern Guaymas Basin. There,  $\text{CH}_4$  generation was related to thermocatalytic degradation of organic matter in sediments intruded by magmatic rocks ( $\text{CH}_4 \sim 6.3 \text{ mM}$ ,  $\text{C1/C2+} \sim 80$ ) (Welhan and Lupton, 1987). The isotopic composition of methane ( $\delta^{13}\text{C} : -39$  to  $-14.9 \text{ ‰}$ ) at our site also indicates thermogenic methane most likely derived from local

sediments, however, with admixture of isotopically heavy methane (Fig. S8). This is consistent with high enrichment in Rn (Tab. S2). The heaviest  $\delta^{13}\text{C}\text{-CH}_4$  is likely related to an abiogenic methane source derived from water rock interaction (e.g., East Pacific Rise-type methane (Welhan and Lupton, 1987)). The presence of abiogenic hydrocarbon is also supported by the isotopic heavy composition of ethane and propane (McDermott et al., 2015; Proskurowski et al., 2008) (Fig. S9). Although the high-temperature aureole at the sill-sediment contact zone appears to be the plausible source for production of hydrocarbons by hydrothermal alteration of organic matter and abiogenic hydrocarbon release by hydrothermal alteration of magmatic rocks (McDermott et al., 2015), our concentration and isotope data do not exclude  $^{13}\text{C}$ -enrichment of methane by (high-temperature) secondary oxidation within the sediments (Pan et al., 2006; Biddle et al., 2012). In fact, the strongest methane input from the smoker vent field exhibits an isotopically light  $\delta^{13}\text{C}$ -trend as it is postulated for the PETM event (Dickens, 2011), and any subsequent methane oxidation product would maintain this signal. We note that isotopically-light carbon of biogenic origin accounts for most of the  $\text{CH}_4$  that is emitted from the black smoker field, despite the presence of an isotopically-heavy hydrocarbon source of abiogenic, magmatic origin. Thus, we conclude that magmatic activity acts mainly as the heat source that fosters and triggers  $\text{CH}_4$  production and release from the post-rift sediments. While abiogenic methane along with MORB-source fluids is indeed emitted, it amounts only to a minor share of the total liberated  $\text{CH}_4$  (mixing ratio:  $1 : 10^2 - 10^4$ , Fig. S8).

## COMPARISON TO MOUND STRUCTURES ON THE NORTH ATLANTIC MARGIN

The Guaymas mound resembles positive relief structures on the Paleocene-Eocene seafloor of the Vøring Basin off mid-Norway that have been interpreted as mud volcanoes previously (Svensen et al., 2004; Aarnes et al., 2015). Although deeply buried, the Norwegian mound structures have similar sizes and internal seismic appearance as the newly discovered active vent structure in the northern Guaymas Basin. The Norwegian mound structures are characterized by a transparent to stratified seismic facies confined by a mostly continuous, low-amplitude seismic reflection (Fig. 2b). The bases of these dome structures also have rugged topography with decreased seismic amplitudes and they overlie zones of disturbed seismic reflections with seismic amplitudes different from their host rocks. These seismic anomalies may indicate sediment alteration during vent activity (Fig. 2b). In addition to similar dimensions, the Norwegian and Guaymas structures both have very steep slopes (Fig. 2c) indicating comparable rheological properties which is uncommon even for the steepest mud volcanoes so far described for marine environments (Pinheiro et al., 2003). Seismic data are never fully conclusive – however, the striking morphological similarity and the location of the structures above seismically imaged fluid pathways indicates that the Guaymas vent may be a suitable analogue for the structures that formed at the beginning of the PETM. This link points to an important role of sediment alteration by ascending hydrothermal fluids and mineral precipitation in the formation of the mounds.

## IMPLICATIONS

Hydrothermal systems injecting hot, CH<sub>4</sub>- and CO<sub>2</sub>-rich fluids high up into the water column are a much more efficient mechanism for releasing large amounts of carbon into the atmosphere than mud volcano-style cold seeps. Such focused ‘hot’ input



efficiently bypasses microbial benthic filters that may oxidize much of the CH<sub>4</sub> to the less potent greenhouse gas CO<sub>2</sub>. Even more important is the observation that hydrothermal systems can inject CH<sub>4</sub> and CO<sub>2</sub> vigorously several hundred meters high up into the water column. In a shallow marine rift environment such as the North Atlantic volcanic rifted margins during the PETM (Planke et al., 2000), such gas plumes may directly reach the atmosphere. Thus, the new observations support the hypothesis that the PETM was caused by the vigorous and wide spread magmatic systems of the North Atlantic large igneous province (Svensen et al., 2004) rather than by large-scale hydrate dissociation.

## ACKNOWLEDGMENTS

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### 313 **FIGURE CAPTIONS**

314 Figure 1. The Guaymas Basin is one of the rift basins formed by opening of the Gulf  
315 of California in NW-SE direction. The recently discovered black smoker is located  
316 just south of the northern rift axis.

317

318 Figure 2. Comparison of the seismic signature of the active black smoker (a) and an  
319 extinct structure in the Guaymas (b and c) and extinct structures observed in the  
320 Vøring Basin during the opening of the Northeast Atlantic off Norway (d and e).  
321 Width (f) and height (g) of the active black smoker in the Guaymas Basin are similar  
322 to structures in the Vøring Basin as indicated by the red line in the blue histograms.  
323 Green lines, average width and height. MTD, mass transport deposit. CSF, chaotic  
324 seismic facies underneath the smoker. Vertical axis on all seismic images shows two  
325 way travel time in ms.

326

327 Figure 3. Three-dimensional view of the mound structure discovered in the Guaymas  
328 Basin. Station IDs of CTD tracks (green lines), HyBis dives (red lines), gravity core  
329 (yellow triangle), and in situ sensor data ( $p\text{CH}_4$  ( $\mu\text{atm}$ ) /  $p\text{CO}_2$  ( $\mu\text{atm}$ ) / temperature  
330 ( $^{\circ}\text{C}$ )) are labelled.

331

332 Figure 4. Schematic diagram illustrating the processes at the active vent site. The  
333 green area indicates under-mature sediments from which organic carbon can be

mobilized by the heat transfer from the magmatic intrusions limiting the depth within which the bulk of the fluids may originate. MTD: Mass transport deposit.

<sup>1</sup>GSA Data Repository item 201Xxxx, including supplementary figures and references as well as additional information on heat flow measurements, geochemical analysis of sediment pore water and water column samples, determination of the sedimentation rate and further seismic information, is available online at [www.geosociety.org/pubs/ft20XX.htm](http://www.geosociety.org/pubs/ft20XX.htm), or on request from [editing@geosociety.org](mailto:editing@geosociety.org) or Documents Secretary, GSA, P.O. Box 9140, Boulder, CO 80301, USA.

#### **GSA Data Repository Materials**

##### ***Seafloor temperature measurements***

The temperature gradients were measured using a 5 m-long temperature-gradient lance with six miniaturized temperature data loggers (MTL). These loggers measured the temperature at a sampling rate of 1 s with an absolute accuracy of approximately 0.1 K (Pfender and Villinger 2002). The relative temperature resolution is 0.001 K. In addition, at six sites a 5 m-long gravity corer with attached MTLs was used. An additional logger at the top of both instruments measured the bottom water temperature as a reference. All measurements were tilt corrected. The derived temperature data were not corrected for sedimentation and terrain effects.

We have measured the thermal conductivity on recovered core material that was sampled at or close to the depth of the measured temperature positions using the KD2 Pro Needle Probe Instrument. For the samples without a thermal-conductivity

measurement, i.e. those without cores, we assumed a constant thermal conductivity of 0.7 W/m K. The data were processed using the method published by Hartmann and Villinger (2002). This method determines undisturbed sediment temperatures from the observed temperature decays.

Heat flow values were calculated by using Fourier's Law as the product of the temperature gradient and thermal conductivity.

We used Bullard method in which the integrated thermal resistance is plotted as a function of temperature. Usually this relationship is linear. However, transient processes such as high sedimentation rates, seasonal temperature fluctuations, focused advection processes or heat generation may cause non-linear deviations. Fourteen out of the 15 presented heat flow sites show a linear relationship between temperatures as function of integrated thermal resistance. Just one site shows non-linear relation at shallow sediment depth.

We interpret the overall high heat flow values with large lateral variations as a result of hydrothermal fluid movements along narrow pathways. The high temperature gradient (Supplementary Table S1) excludes conductive heat transfer. Conductive heat transfer cannot explain either the large lateral heat flow variations on a spatial scale of 100 m. This indicates that there is rigorous hydrothermal venting. All of the SO241-70 sites show advective components. Three out of these four are interpretable as discharge areas (in the northern and central part of the graben), but the Bullard plot of (Site SO241-70P04) shows a downward concave curve at shallow depth, which may be the result of seawater recharge into the crust before it is being heated up.

**Geochemistry**

Water samples were taken by using a towed SBE9-CTD rosette device equipped with 11 Niskin bottles (a 10 l) and additional HydroC-pCH<sub>4</sub> and -pCO<sub>2</sub>, turbidity, and bottom distance sensors (modified after Schmidt et al., 2015). CTD-Stations VCTD09 (and -10) were towed directly above the hydrothermal vent field in variable distance to seafloor (Fig. 3). However, only VCTD09 data is shown in Table S2 as it exhibit notable inorganic variations of hydrothermal tracers in water masses in 5-20 m distance to seafloor. Niskin bottles were closed when temperature, pCO<sub>2</sub> and turbidity suddenly increased indicating hydrothermal plume anomalies. The Niskin bottles were sub-sampled directly after CTD retrieval for inorganic element chemistry, nutrients, partial pressures of dissolved gases and stable and radio-isotope characteristics (Tab. S2) and water samples were analysed onboard for nutrient concentrations. Sub-samples taken for the analysis of major cations (Na<sup>+</sup>, K<sup>+</sup>, Li<sup>+</sup>, Mg<sup>2+</sup>, Ca<sup>2+</sup>, Sr<sup>2+</sup>, Ba<sup>2+</sup>) as well as B and Si were measured in the shore-based laboratory at GEOMAR Helmholtz-Centre for Ocean Research by ICP-OES (<http://www.geomar.de/en/research/fb2/fb2-mg/benthic-biogeochemistry/mg-analytik/icp-aes/>). ICP sub-samples were acidified directly after sampling to prevent any precipitation of minerals. The elements Ba, Mn, Si, and Li are enriched compared to seawater values. The concentration ranges between 128 – 1766 nM for Ba, 0.02 to 23.9 µM for Mn, 0.16 – 0.69 mM for Si and 23.1 – 59.2 µM for Li (Table S2). All elements show the strongest anomalies (mostly enrichment) in bottle 12 (VCTD09). Mg shows a slight concentration decrease compared to seawater. The strongest depletion of Mg (51 mM) was also encountered in bottle 12. TIMS measured <sup>87</sup>Sr/<sup>86</sup>Sr ratios normalized on SRM-987 (0.710248) ranges between 0.708906 and almost

modern seawater (0.709176) matching 0.709170 ( $\pm 1 \text{ E-5}$ , typical 2 SEM this session) showing the lowest value in bottle 12 which was sampled closest to the active vent.

We calculated the amount of hydrothermal fluids in the Guaymas Basin bottom water using a simple two-end member mixing model between the seawater and primary hydrothermal fluid which was assumed to be void of Mg. Based on this we calculated the maximum percentage of the hydrothermal fluid in the water samples. The lowest Mg concentration of 51 mM in the water sample of bottle 12 yields a hydrothermal fluid percentage of ~6 %. Based on this we determined the hydrothermal end member composition of the enriched elements in the sampled water column and compared those to the concentrations measured at the southern Guaymas spreading center by Von Damm et al. (1985).

Hydrothermal end member concentrations for Si and Ba yielded 9.36 mM and 28  $\mu\text{M}$ , respectively and fall well into the end member concentration ranges observed by Von Damm et al. (1985) in the southern Guaymas Basin (Si: 9 - 14 mM; Ba: 7 – 42  $\mu\text{M}$ ). Si concentration is at the lower end of the concentration range and might point to a lower hydrothermal temperature or Si precipitation during ascent or after fluid discharge. Hydrothermal end member concentrations calculated for Mn yield 420  $\mu\text{M}$ , which is higher than the concentrations observed in the southern Guaymas hydrothermal fluids, which range between 128 and 236  $\mu\text{M}$  (Von Damm et al., 1985). Von Damm et al. (1985) propose the precipitation of alabandite (MnS) as an explanation for the observed low values. Li, in contrast, shows slightly lower calculated end member concentrations (584  $\mu\text{M}$ ) compared to the southern Guaymas fluids, which range between 630 and 1076  $\mu\text{M}$ . These lower values might be



explained by a Li sink in the sediments through which the hydrothermal fluids percolate.

Concentrations of dissolved He and Ne (as well as Ar, Kr and Xe) and the  $^3\text{He}/^4\text{He}$ ,  $^{22}\text{Ne}/^{20}\text{Ne}$ ,  $^{36}\text{Ar}/^{40}\text{Ar}$  isotope ratios in water samples taken in copper tubes were analysed as routine samples in the noble gas laboratory of the Swiss Federal Institute of Technology Zurich (ETHZ) and the Swiss Federal Institute of Aquatic Science and Technology (Eawag). Details on sampling and of the experimental method are given in Beyerle et al. (2000). He and Ne concentrations are covered with an overall standard error of  $\pm 2\%$ , the  $^3\text{He}/^4\text{He}$  ratio has a standard error of 1%.

For the on-board analysis concentrations of dissolved He, Ar, Kr,  $\text{N}_2$ ,  $\text{O}_2$ ,  $\text{CO}_2$ , and  $\text{CH}_4$  during the expedition, we used a portable gas-equilibrium membrane-inlet mass-spectrometric system (GE-MIMS) similar to that described in Mächler et al. (2012). Whereas the original GE-MIMS was designed for gas analysis in continuous high-volume water flows through a large membrane contactor, the GE-MIMS used here was modified to allow analysis of an 8 l water sample taken from a 10 l Niskin sampler within 10 min (Brennwald et. al., 2015, a, b). To allow reliable gas analysis in this limited amount of water, the gas consumption from the membrane contactor into of the MS was reduced to 0.1 ccSTP/min ( $> 2$  ccSTP/min in the original GE-MIMS). This allowed maintaining gas/water solubility equilibrium in miniature membrane contactor modules (two LiquiCel MicroModules operated in parallel) at a total water throughput of 0.5 L/min ( $> 5$  L/min in the original GE-MIMS). The low gas consumption was achieved by replacing the capillary flow resistance followed by a split-flow/aperture gas inlet to the MS vacuum by a new splitless design using a

single capillary (9 m long, 0.1 mm inner diameter, Brennwald et al., 2015b). The gas outflow from the capillary is analysed in a quadrupole MS (Stanford Research Systems RGA 200) operated in dynamic mode. As the membrane contactors operate at solubility equilibrium, the partial pressures of the noble gases N<sub>2</sub> and O<sub>2</sub> in the gas phase of the membrane contactors are similar to those in air. The GE-MIMS data for these species were therefore calibrated using ambient air as a reference gas (< 5 % accuracy, 1  $\sigma$ ). The partial pressures of CO<sub>2</sub> and CH<sub>4</sub> are reported as un-calibrated raw data.

In order to determine the variation of excess amounts of the noble gas radon ( $^{222}\text{Rn}_{(\text{exc.})}$ ) close to the hydrothermal vent, selected CTD samples were measured on board by Liquid Scintillation Counting (LSC). Using a portable Hidex Triathler<sup>TM</sup> system and MaxiLight<sup>TM</sup> as scintillation cocktail for the measurement of total  $^{222}\text{Rn}$  content uncertainties of 15% (SD) are typical for the counting statistics. After more than 3 months storage re-measurements were conducted at GEOMAR to correct for the fraction of total  $^{222}\text{Rn}$  potentially provided by the decay of dissolved  $^{226}\text{Ra}$ . The analytical procedure followed the approach described by Purkl and Eisenhauer (2004). The maximum sample size was restricted to 1.5 l combined with 20 ml of LSC cocktail, which is close to the maximum extraction efficiency as recently deduced by Schubert et al. (2014). The re-measurements for supported  $^{222}\text{Rn}$  contribution did not reveal concentrations above the detection limit of the applied LSC setup. Consequently, a simple, robust, and reliable semi-quantitative measure on the relative enrichment in  $^{222}\text{Rn}_{(\text{exc.})}$  can be presented in Tab. S2 in terms of enrichment factors compared to bottle 5. Since bottle 5 shows the lowest  $^{222}\text{Rn}$  activity in dpm/L (decay per minute/liter) of the investigated CTD casts it provides

the best available approximation to water column background  $^{222}\text{Rn}$  activity and was set as reference point (Tab. S2). The lowest published  $^{222}\text{Rn}$  data (0.1 and 0.2 dpm/L) of Santos et al. (2011) for central waters of the Concepcion Bay (West of our study area) and of Prol-Ledesma et al. (2013) for the northern Gulf of California (Wagner and Consag basins, down to zero dpm/100L) may be assumed as typical background values and are similar to the lowest values that we have encountered (bottle 5, 1768 m: 0.17 dpm/L ( $\pm 0.1$ , SD; calibration pending). However, applying the same approach to the sample recovered closest to the vent (bottle 12, 1773 m) yielded the maximum  $^{222}\text{Rn}_{(\text{exc.})}$  activity of 13.8 dpm/L for our data set. Independent from exact quantitative setup calibrations, this approach provides a robust enrichment factor for  $^{222}\text{Rn}_{(\text{exc.})}$  in the same order of magnitude as deduced for He in this study (Tab. S2). However, almost twice as high levels of  $^{222}\text{Rn}_{(\text{exc.})}$  of 2430 dpm/100L were reported for the hydrothermal impact on the waters of the northern Gulf of California in the Wagner and Consag basins (Prol-Ledema et al., 2013). There they attribute the positive anomalies of  $^{222}\text{Rn}$  to a fault system coinciding with the presence of strong flares suggesting hydrothermal circulation in a thick sediment cover and revealing the location of up-flow areas.

Dissolved hydrocarbons (C1-C3) from individual water samples were released onboard by equilibration of 112 ml water samples in a septum-sealed 117 ml headspace vial at room temperature (He-head space, 50  $\mu\text{l}$   $\text{HgCl}_2$ -solution added). Hydrocarbon composition of the head space gas was determined by using a Thermo Trace gas chromatograph (GC) equipped with flame ionization detector (carrier gas: He 5.0; capillary column: RT Alumina Bond-KCl, column length: 50 m; column diameter: 0.53  $\mu\text{m}$ ). Precision of  $\pm 1$ -3% was achieved when measuring standard

hydrocarbon mixtures. Molar hydrocarbon concentrations in Table S2 were calculated by applying Henry coefficients according to Mackay and Shiu (2006).

Stable carbon isotope ratios of methane and higher hydrocarbons (C1-C3) from water samples were measured by using continuous flow GC combustion - Isotope Ratio Mass Spectrometry. Hydrocarbons were separated in a Thermo Trace GC (carrier gas: He; packed column: ShinCarbon, 1.5 m). The subsequent conversion of hydrocarbons to carbon dioxide was conducted in a Ni/Pt combustion furnace at 1150°C. The  $^{13}\text{C}/^{12}\text{C}$ -ratios of the produced  $\text{CO}_2$  were determined by a Thermo MAT253 isotope ratio mass spectrometer. All isotope ratios are reported in the  $\delta$ -notation with respect to Vienna Pee Dee Belemnite (VPDB, analytical precision 0.5 ‰).

### ***Sedimentation Rates***

The wet sediment was weighed, freeze dried at -80°C and reweighed to determine the water content, then ground in a mortar. Radionuclides were then measured as follows. Two HPGe detectors were engaged for  $^{210}\text{Pb}$  and  $^{226}\text{Ra}$  analysis including GMX-type (ORTEC GMX-120265) and well-type (ORTEC GWL-100230) detectors which interfaced to a digital gamma-ray spectrometer (DSpecPlus™). For the GMX-type detector, absolute counting efficiencies for various photon energies were calibrated using IAEA reference materials 327A, 444 spiked soil, CU-2006-03 spiked soil, RGTh and RGU for sample weight at 100g as a reference, and coupled with an in-house secondary standard for various masses (from 10 to 250 g) to calibrate the effect of sample mass on the attenuation of  $\gamma$ -rays of various energies. For the well-type detector, the counting efficiencies were calibrated by IAEA-RGTh and RGU from 0.5 to 3.5 g.  $^{214}\text{Pb}$  was used as an index of  $^{226}\text{Ra}$  (supported  $^{210}\text{Pb}$ ) whose activity

concentration was subtracted from that of the measured total  $^{210}\text{Pb}$  to obtain excess  $^{210}\text{Pb}$  ( $^{210}\text{Pb}_{\text{ex}}$ ). The  $^{210}\text{Pb}$  and  $^{214}\text{Pb}$  activities were quantified based on photon peaks centered at 46.52 and 351.99 keV, respectively. The activities of radionuclides were decay-corrected to the date of sample collection. All radionuclide data were calculated on salt-free dry weight basis. Error bars represent  $\pm 1\sigma$  around the mean based on counting statistics and standard propagation of errors.

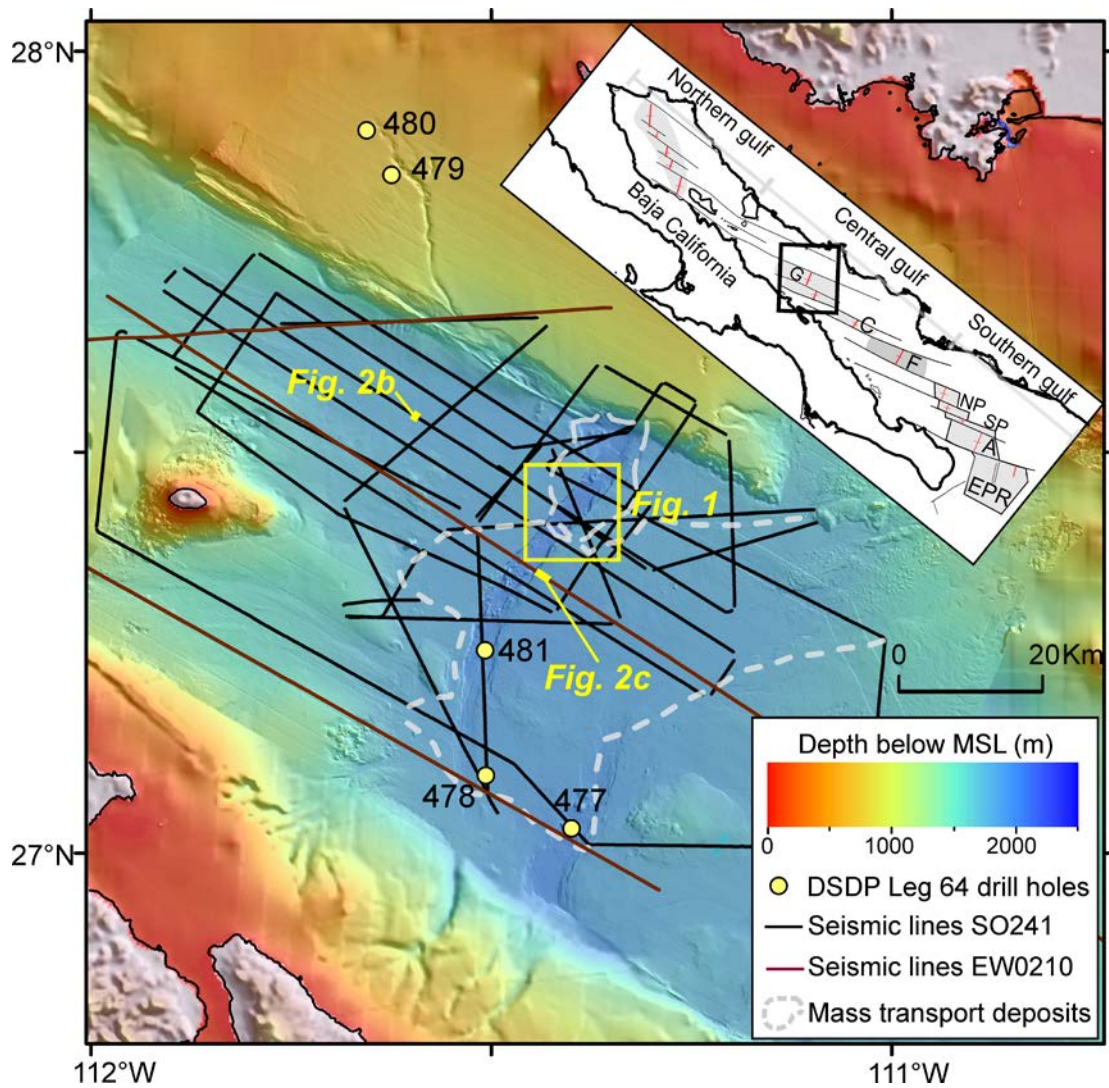
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593 Figure S1: Seismic base map showing the distribution of available 2D seismic data in  
 594 the Guaymas Basin and major structural elements (after Padilla y Sánchez et al.,  
 595 2013). Central and Southern gulf spreading centers are labelled, G–Guaymas, C–  
 596 Carmen, F–Farallon, NP–North Pescadero, SP–South Pescadero, A–Alarcon, EPR–  
 597 East Pacific Rise.

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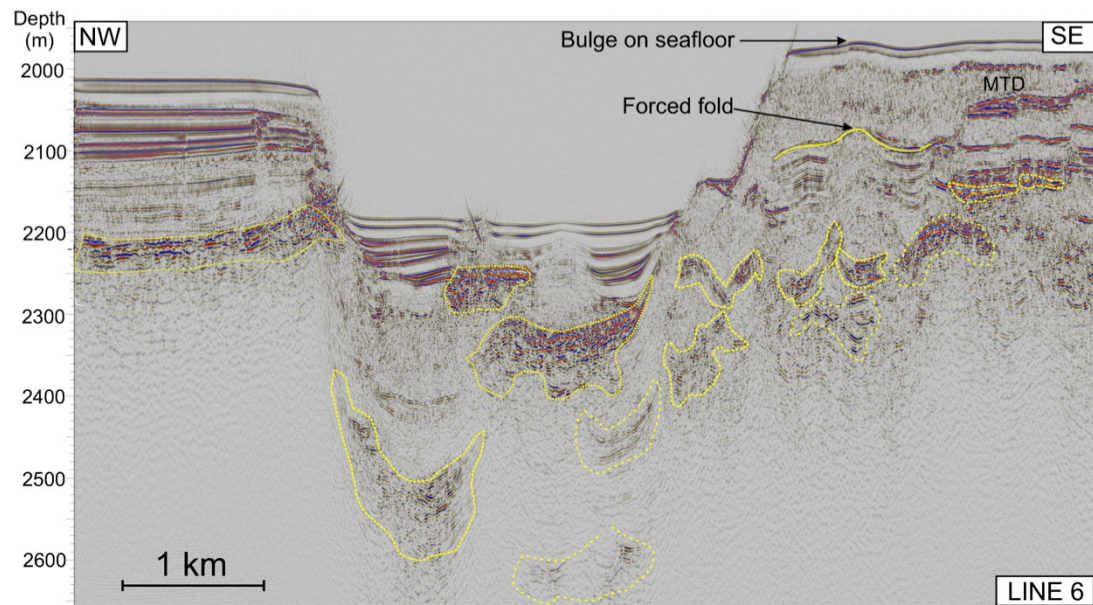
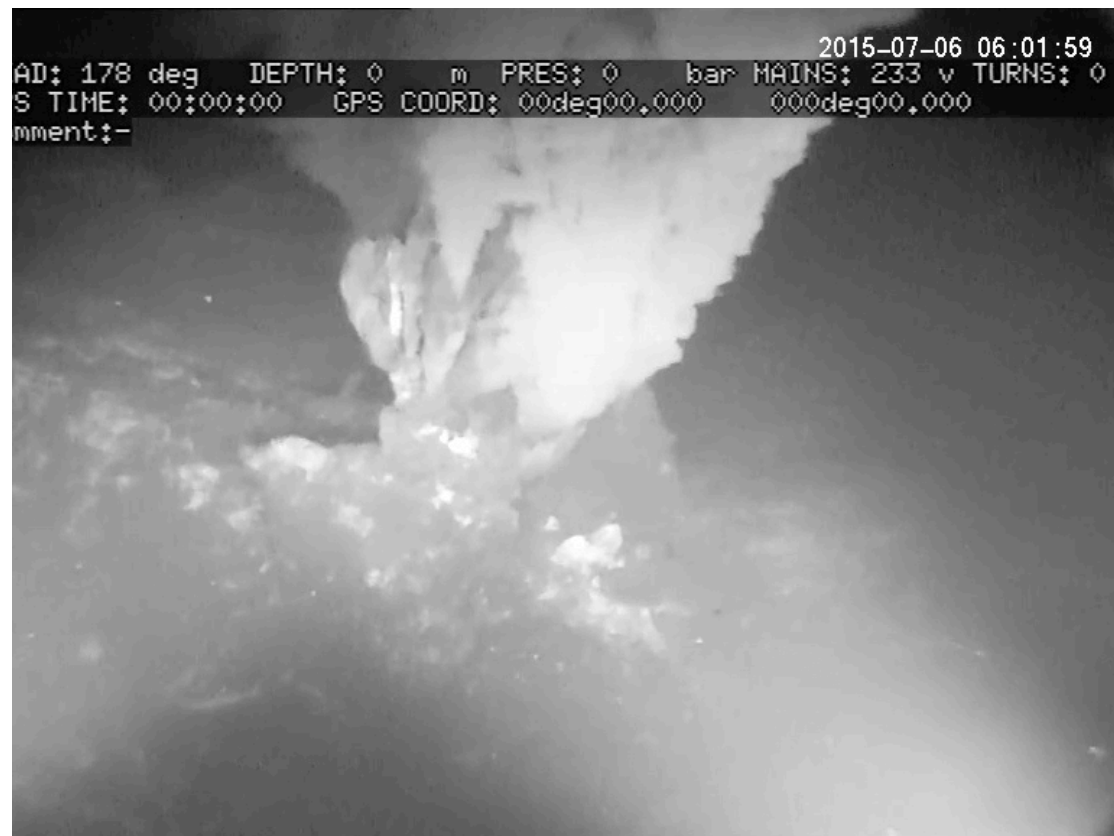


Figure S2: 2D seismic line showing the regional mass transport deposit (MTD) and interpreted sill intrusions (yellow polygons).



Figure S3: Photograph of a massive sulfide sample collected on the southern summit of the mound structure.



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611 Figure S4: Video still of one of the seven active vents on top of the mound structure.

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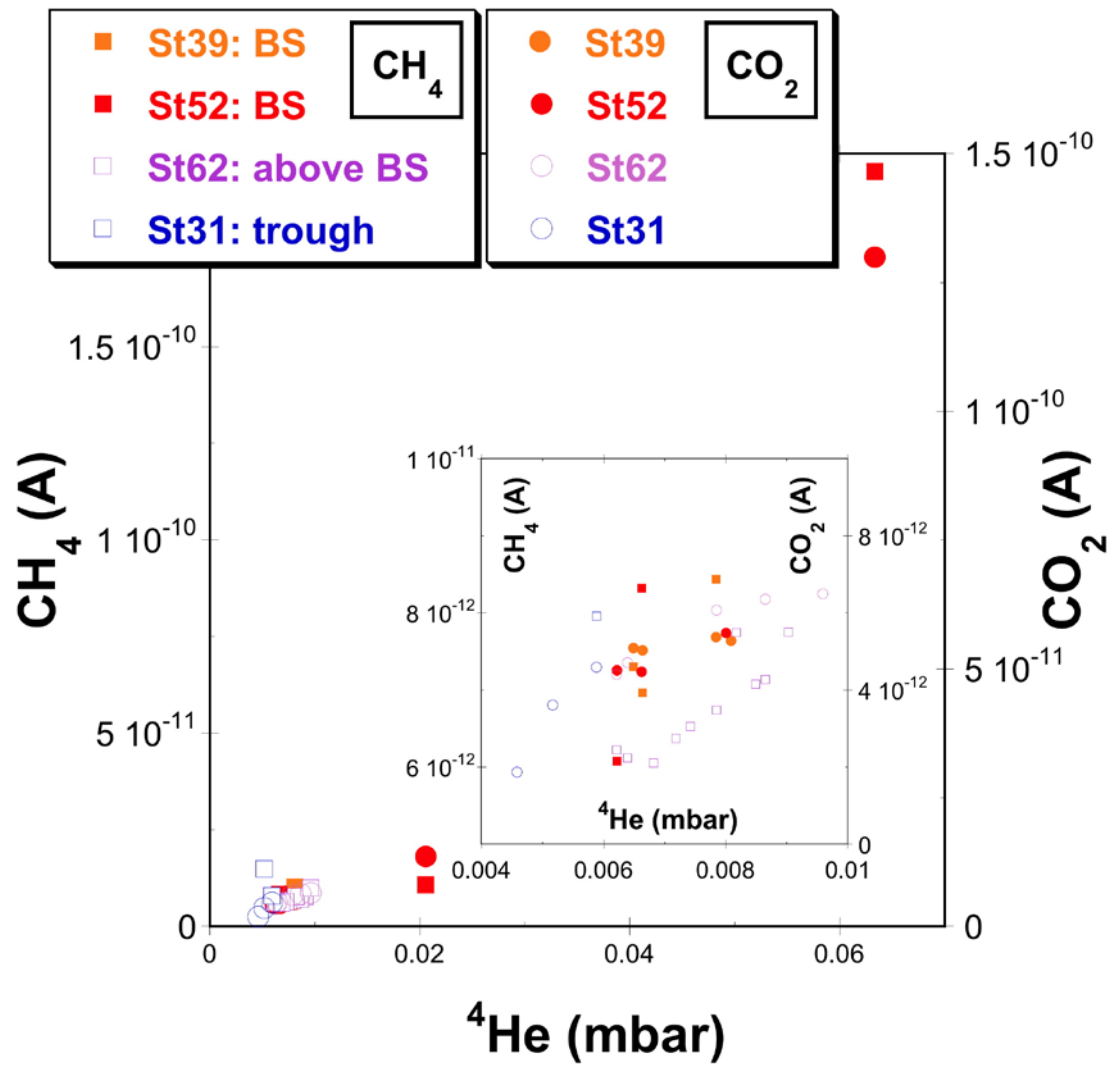


Figure S5: MIMS-determination of dissolved  $^4\text{He}$ ,  $\text{CH}_4$  and  $\text{CO}_2$  partial pressures (mbar) measured at, above and in the vicinity of the hydrothermal vent site (Stxy: station number xy, BS: 'Black Smoker - within the hydrothermal field (stations: 39, 52)', above BS: 'free water column above vent sites (station 62)', trough: samples in the deep trough of the northern Guaymas Basin (station 31)).

The partial pressures of  $\text{CO}_2$  and  $\text{CH}_4$  are reported as un-calibrated raw MIMS detector signals.

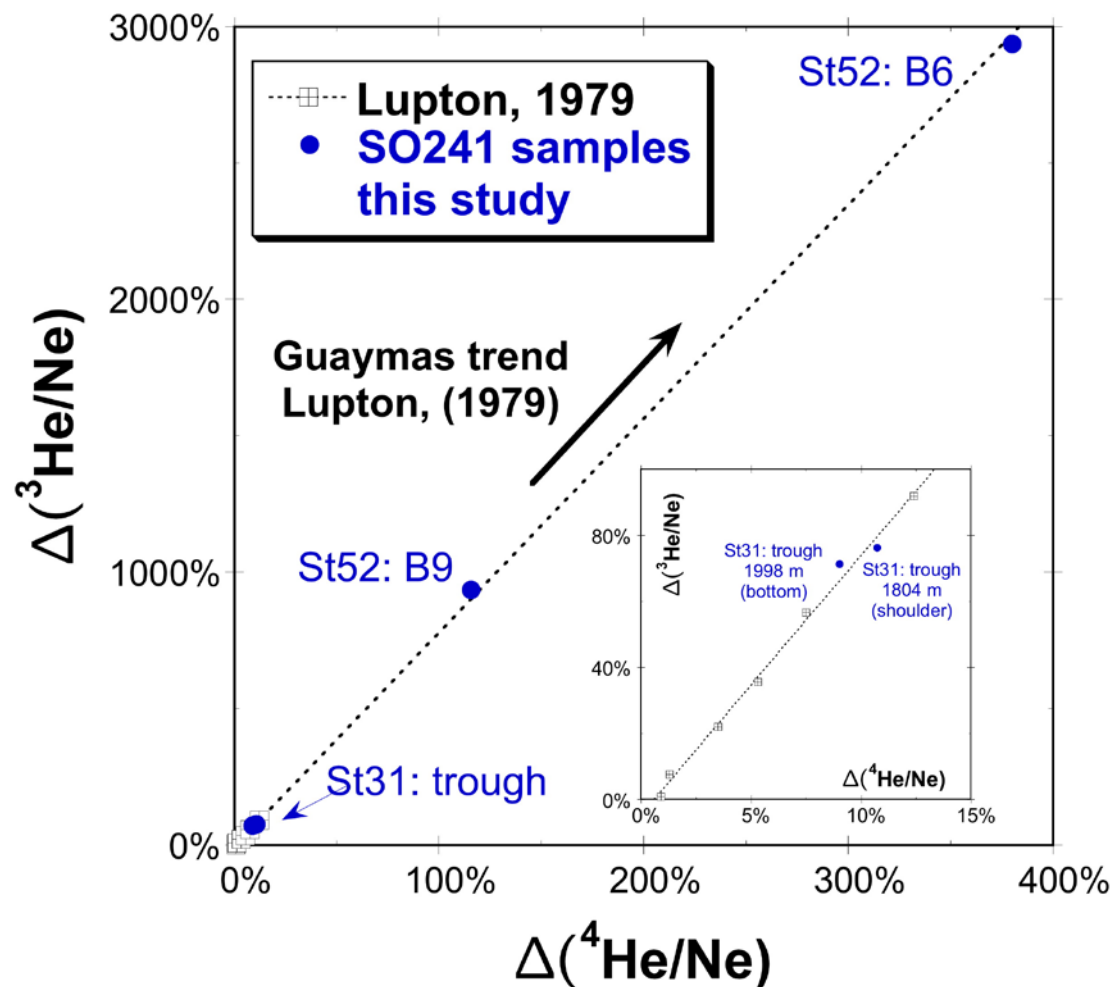


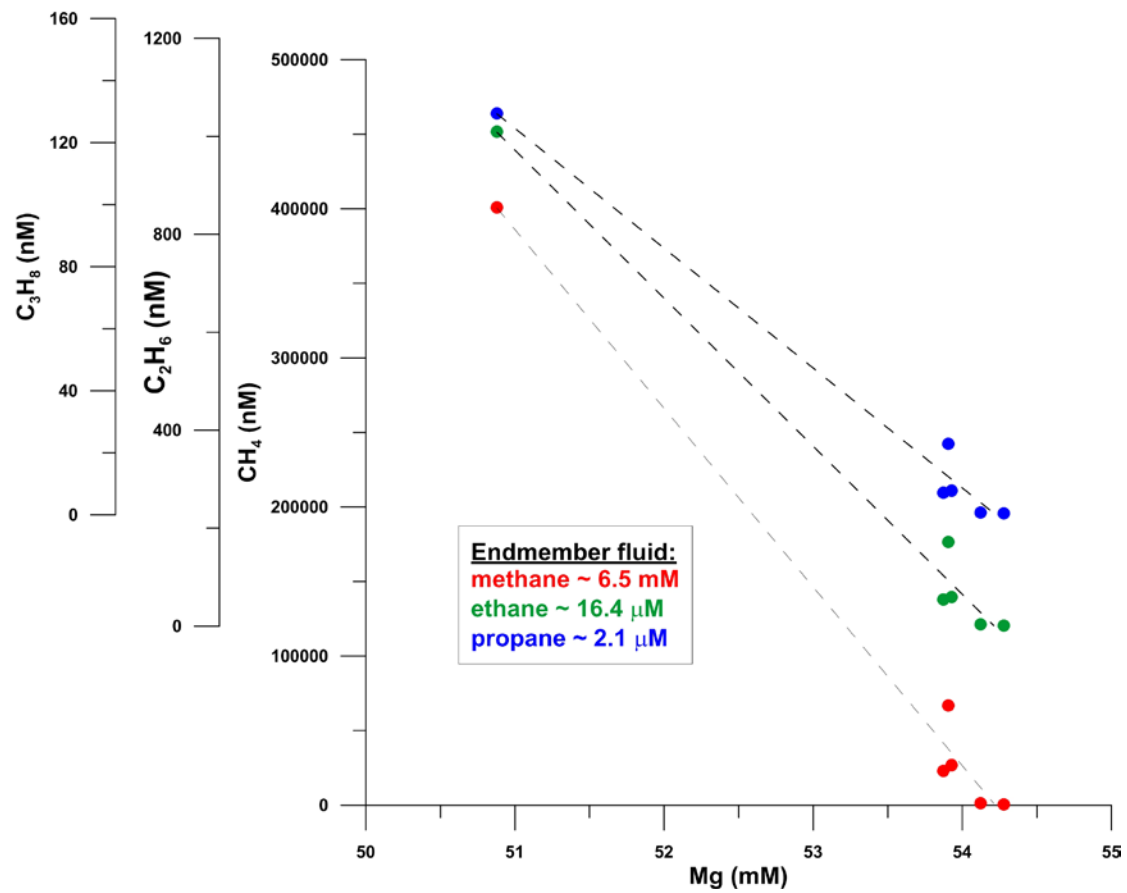
Figure S6: Dissolved He and Ne concentrations measured at the hydrothermal vent site in comparison to earlier noble gas measurements in the southern part of the Guaymas Basin (Lupton, 1979). The samples from the vent site fall on the line defined by the earlier measurements from the southern Guaymas Basin and thus carry isotopic light He ( $^3\text{He}/^4\text{He}$ :  $10.8 \cdot 10^{-6}$ , Lupton, 1979) from the same general MORB source. For comparison noble gas (isotope) concentrations are reported and normalized as in Lupton, 1979:

$$\Delta\left(\frac{^i\text{He}}{\text{Ne}}\right) = \left(\frac{^i\text{He}_m}{\text{Ne}_m} \times \frac{\text{He}_{\text{ASW}}}{\text{Ne}_{\text{ASW}}} - 1\right) \times 100 \%$$

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634 where  $X_m$  depicts the measured concentration of X, and  $X_{ASW}$  is the expected  
635 atmospheric equilibrium concentration (atmospheric saturated water: ASW) for the  
636 given physical condition of the water. As  $^i\text{He}_{ASW}$  and  $\text{Ne}_{ASW}$  only barely change with  
637 varying temperature and salinity the  $\text{Ne}_{ASW}/^i\text{He}_{ASW}$  ratio is virtually independent of  
638 the actual physical condition of the water. Stxy: 'station number, 'Bz': bottle number.  
639 Station 52: Black Smoker site, Station 31: open water column of the trough.

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643 Figure S7: Hydrocarbon end member concentrations (methane, ethane, propane)

644 calculated for hydrothermal fluids venting at the smoker area.

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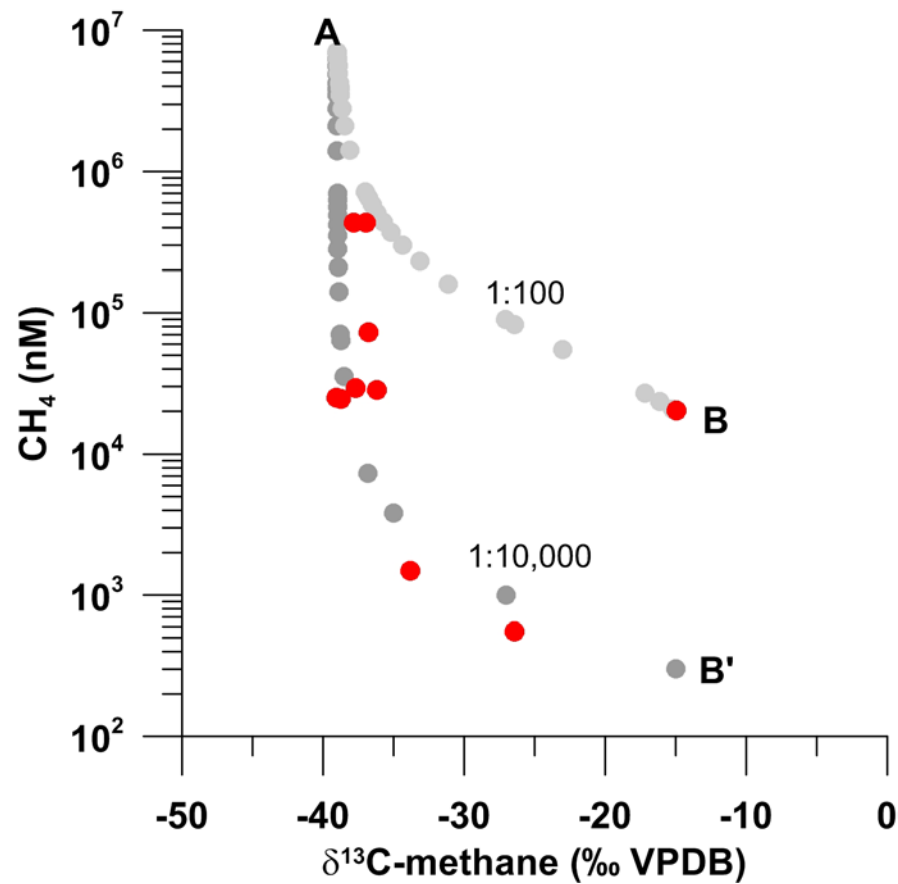


Figure S8: Dissolved methane concentration and carbon isotope data (red dots) determined in bottom water samples from the smoker area. Grey dots indicate mixing curves of a hydrothermal end member methane A ( $\delta^{13}\text{C} = -39$  ‰, 7nM) with admixture of a second methane-containing fluid B and B' ( $\delta^{13}\text{C} = -15$  ‰), respectively, at variable dilution factors of 1:100 and 1:10,000 compared to fluid A.

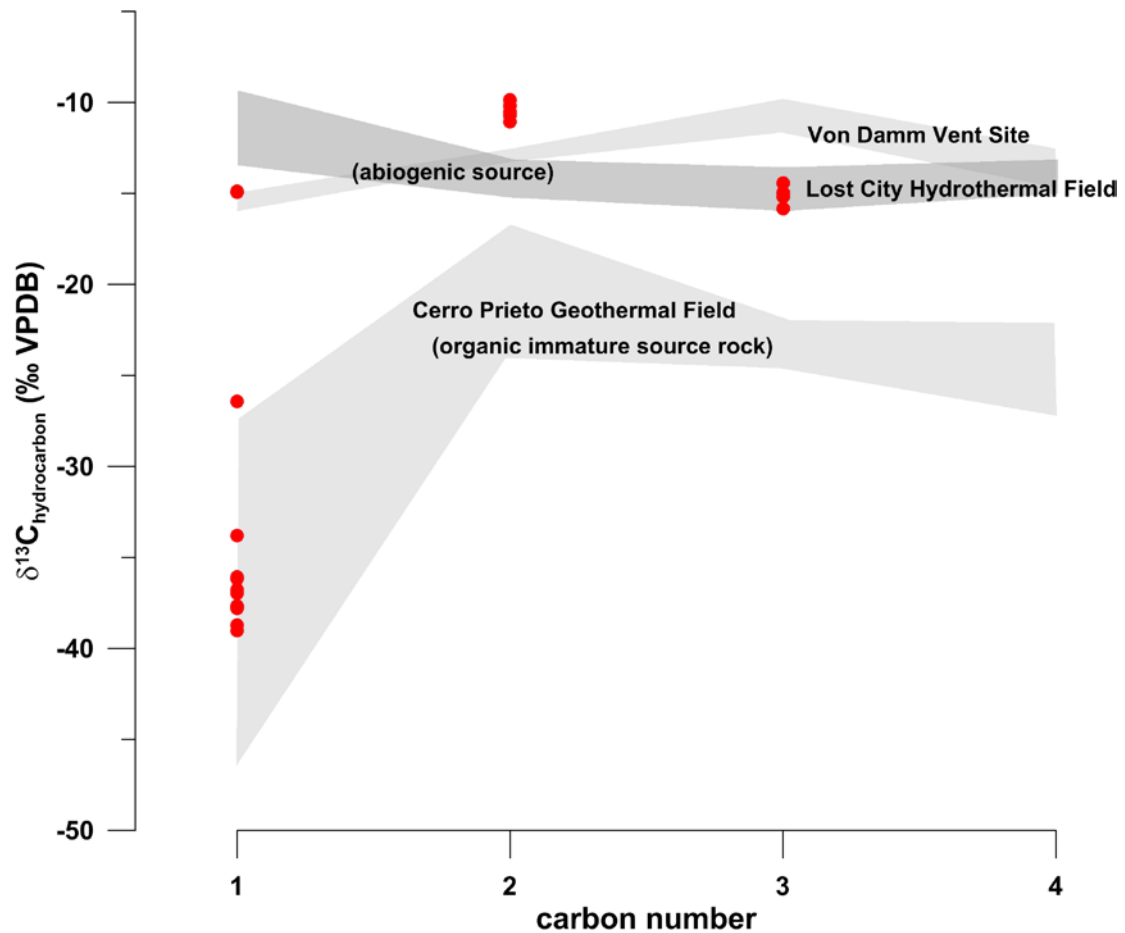


Figure S9: Stable carbon isotope composition of hydrocarbons dissolved in venting hydrothermal fluids (red dots) indicate both hydrocarbons derived from organic matter degradation influenced by volcanic heat intrusion (e.g. Cerro Prieto volcanic complex; Des Marais, 1988), and abiogenic hydrocarbon formation (e.g. Lost City HF, Proskurowski et al., 2008; Von Damm Vent, McDermott et al., 2015).



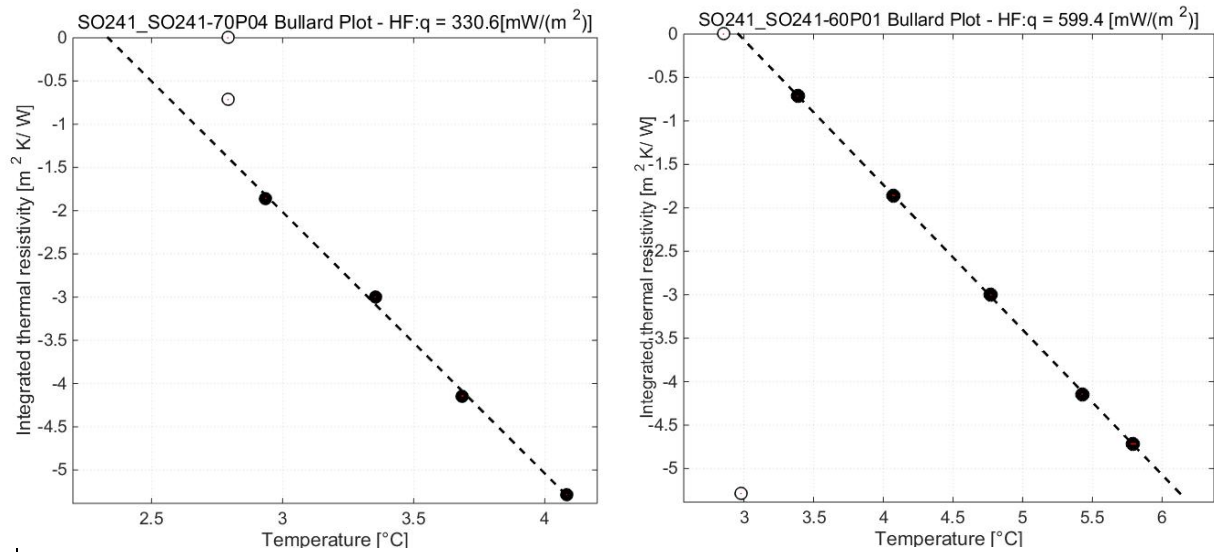


Figure S10: Calculated heat flow for Site SO241-70P04 (top) and SO241-60P01 (bottom) using the Bullard Plot method. Black dots represent sensors used to calculate the heat flow whereas unfilled dots represent the sensor positions, which were not used for any calculations because they were affected by surface artifacts such as incomplete penetration of the lance.

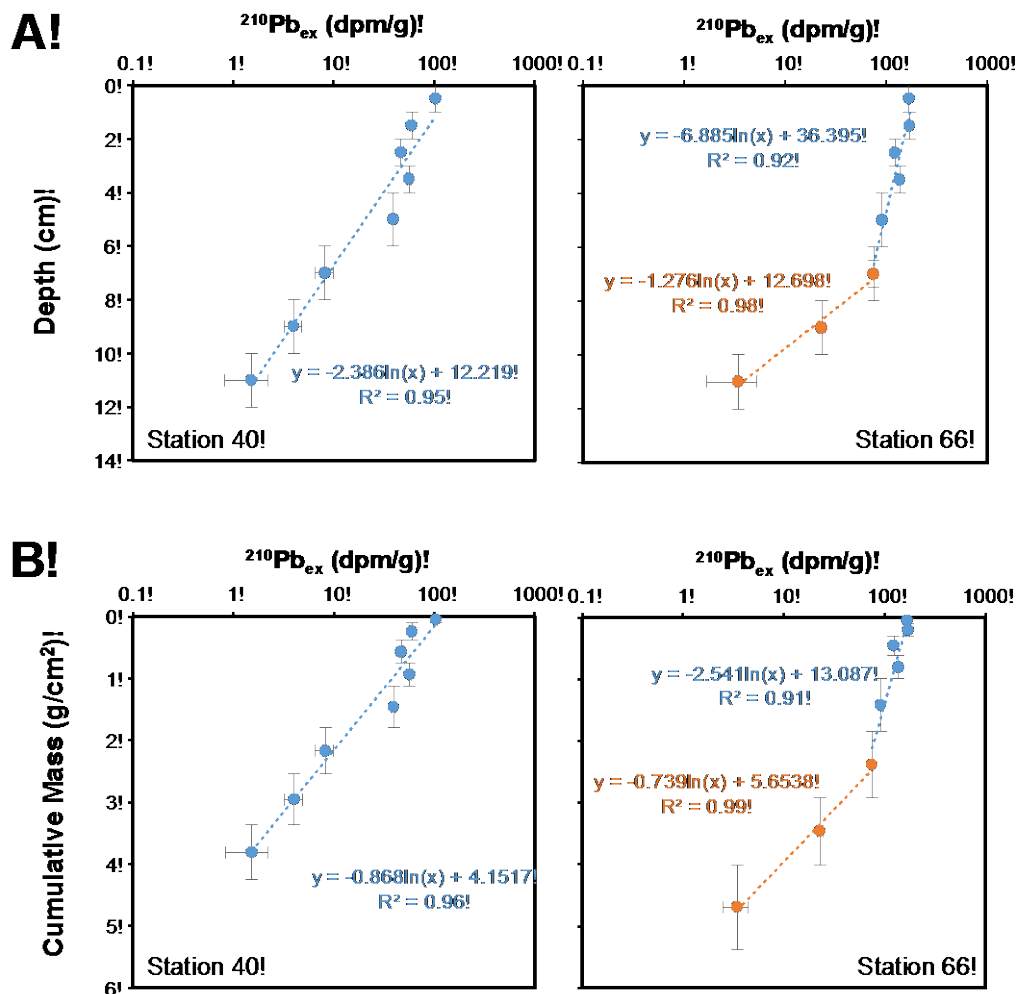


Figure S11:  $^{210}\text{Pb}_{\text{ex}}$  values plotted versus (A) sediment depth and (B) cumulative mass at Stations 40 and 66. Lines and equations represent linear-best fit functions and their data fit ( $R^2$ ). Station 40 has a good linear fit from the surface to the deepest layers sampled, indicating a relatively steady sedimentation rate over the sampling interval. By contrast, Station 66 has two best-fit lines, one to match samples from the top 6 cm (cyan), and a second one to match samples from 6-12 cm (orange). Samples from the top 6 cm at Station 66 show an only minor decrease in  $^{210}\text{Pb}_{\text{ex}}$ , presumably due to vertical sediment mixing by macrofauna. The samples from 6-12 cm at Station 66 show a stronger  $^{210}\text{Pb}_{\text{ex}}$  decrease with depth that more likely reflect the sedimentation rate of the site and were thus used for calculations of sedimentation

681 rates. While the profiles of  $^{210}\text{Pb}_{\text{ex}}$  show good agreement independent of the y-axis  
682 units at Station 40, there is a considerable difference between  $^{210}\text{Pb}_{\text{ex}}$  profiles at  
683 Station 66 depending on whether sediment depth or cumulative mass are used as y-  
684 axis units. This discrepancy is likely an artefact of core compaction during sediment  
685 sampling. We consider  $^{210}\text{Pb}_{\text{ex}}$  relationships with cumulative mass, which correct for  
686 compaction effects during coring, to be more reliable (especially at Station 66), and  
687 thus report sedimentation rates that have been corrected for cumulative mass in this  
688 article.

**Supplementary Tables**

Table S1: Heat flow measurements carried out during SO241

Station	Penetration	Latitude	Longitude	Water Depth [m]	Temp. Gradient [K/m]	k [W/mK]	Heat flow [mW/m <sup>2</sup> ]
SO241-51	Pen 01	27° 24.472'	111° 23.377'	1840	11.441	0.739	8069.3
SO241-58	Pen 01	27° 24.487'	111° 23.377'	1837	9.857	0.720	6508.7
SO241-60a	Pen 01	27° 24.623'	111° 23.626'	1834	0.856	0.7 *	599.4
	Pen 02	27° 24.554'	111° 23.512'	1840	2.789	0.7 *	1952.6
	Pen 03	27° 24.273'	111° 23.396'	1840	4.581	0.7 *	3206.1
	Pen 04	27° 24.408'	111° 23.288'	1849	2.039	0.7 *	1427.0
	Pen 05	27° 24.341'	111° 23.177'	1852	1.014	0.7 *	709.6
	Pen 06	27° 24.265'	111° 23.082'	1844	0.737	0.7 *	516.1
	Pen 07	27° 24.193'	111° 22.956'	1834	0.827	0.7 *	578.7
SO241-60b	Pen 01	27° 24.605'	111° 23.317'	1837	0.391	0.7 *	274.0
	Pen 02	27° 24.552'	111° 23.347'	1834	3.451	0.7 *	2415.4
	Pen 04	27° 24.543'	111° 23.351'	1837	15.479	0.7 *	10835.0
SO241-70	Pen01	27° 25.802'	111° 25.486'	1870	0.375	0.7 *	262.2
	Pen02	27° 25.460'	111° 24.946'	2019	0.483	0.7 *	338.1
	Pen03	27° 25.955'	111° 24.493'	2046	0.432	0.7 *	302.6
	Pen 04	27° 24.837'	111° 23.951	2025	0.457	0.7 *	319.7

Thermal conductivities k with an asterisk (\*) are assumed.

Bottle	Latitude	Longitude	Depth	Temperature	Salinity	Methane (C1)	Ethane (C2)	Propane (C3)	δ <sup>13</sup> C-C1	δ <sup>13</sup> C-C2	δ <sup>13</sup> C-C3	<sup>222</sup> Rn <sub>(exc.)</sub> <sup>a</sup>	Mn	Mg	Sr	<sup>87</sup> Sr/ <sup>86</sup> Sr	Si	Ba	Li
	N	W	(m)	(°C)	(‰)	(nM)	(nM)	(nM)	(‰ VPDB)	(‰ VPDB)	(‰ VPDB)	enrich. fact.	(μmol/l)	(mmol/l)	(μmol/l)		(mmol/l)	(nmol/l)	(μmol/l)
1	27.41258	-111.3870	1504	3.1	34.60														
2	27.41284	-111.38696	1766	2.9	34.61	1366.1	3.7	0.8	-33.8				0.02	53.0	88.6	0.70917	0.16	128	23.1
3	27.41288	-111.38696	1767	2.9	34.61							1.7	<d.l.	54.1	88.7	0	0.18	145	24.9
4	27.41204	-111.38732	1768	4.1	34.55	508.2	0.8	0.4	-26.4				0.14	52.8	88.2	0.70916	0.18	147	23.3
5	27.41204	-111.38732	1768	4.1	34.64							1.0	<d.l.	54.3	89.2	7	0.18	143	24.9
6	27.4121	-111.38732	1768	4.3	34.68	66885.7	171.7	22.9	-36.8	-9.9	-15.8		0.49	52.9	88.4	0.70912	0.18	451	23.8
8	27.4121	-111.38732	1768	4.4	34.62							16.2	3.79	53.9	89.6	9	0.26	946	29.0
9	27.41212	-111.38734	1772	6.9	34.64	26801.6	59.1	7.8	-37.7	-10.5			1.33	53.9	88.6	0.70915	0.20	410	26.5
						26010.5	57.4	8.2	-36.2							6			
						22987.6	54.2	7.1	-39.0	-11.1						0.70915			
10	27.41214	-111.3872	1775	17.4	34.96	22466.2	52.1	6.8	-38.7			4.5	1.18	53.9	88.9	9	0.20	383	26.3
									-14.9										
11	27.41214	-111.3872	1775	28.0	34.17	18684.1	42.6	6.1	-14.9	-10.7	-14.4		1.10 <sup>b</sup>	52.9 <sup>b</sup>	88.3 <sup>b</sup>		0.19 <sup>b</sup>	353 <sup>b</sup>	24.7 <sup>b</sup>
									-14.9										
						360284.0	849.3	137.4		-10.2	-15.0								
12	27.41212	-111.38718	1773	12.1	34.26		893.0	110.2	-37.0			81.5	23.9	51.0	90.6	0.70890	0.69	1766	59.2
						400801.8	1009.3	129.4	-37.8							6			

<sup>a</sup>Enrichment factor of <sup>222</sup>Rn relative to the lowest value measured on this CTD station (bottle 5). See text for details.

<sup>b</sup>Inorganic geochemistry values are a mixture of bottle 10 (~20%) and bottle 11 (~80%)

695 Table S3: Isotope ratios of He and Ne reported as the percentage deviations of <sup>3</sup>He/Ne and  
696 <sup>4</sup>He/Ne from the solubility ratios (see Lupton (1979) for details).  
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Location	Bottle	Latitude	Longitude	$\Delta(^3\text{He/Ne})$ [%]	$\Delta(^4\text{He/Ne})$ [%]
		N	W		
Black Smoker	6	27.4121	-111.38732	2937	380
Black Smoker	9	27.41212	-111.38734	936	116
Background trough bottom	2	37.30207	-111.52433	71	9
Background trough shoulder	7	37.30207	-111.52433	76	11

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